- 14. The method according to claim 9, wherein [the] at least one anti-LT-β-R monoclonal antibody [ies are] is CBE11 and at least one anti-LT-β-R monoclonal antibody is BHA10.
- 15. The method according to claim 9, wherein [the] <u>at least one</u> anti-LT-β-R monoclonal antibody [ies are] <u>is</u> CBE11 and <u>at least one anti-LT-β-R monoclonal antibody is</u> CDH10.
- 16. The method according to claim 9, wherein [the] at least one anti-LT-β-R monoclonal antibody [ies are] is AGH1 and at least one anti-LT-β-R monoclonal antibody is CDH10.
- 17. The method according to any one of claims 6-16, [wherein one LT- β -R activating agent is] <u>further comprising IFN- γ </u>.
- 38. A pharmaceutical composition comprising a therapeutically effective amount of at least two LT-β-R activating agents, and a pharmaceutically acceptable carrier

 [The pharmaceutical composition according to claim 37], wherein at least one LT-β-R activating agent comprises an anti- LT-β-R antibody.
- 46. The pharmaceutical composition according to claim 41, wherein at least one [the] anti- LT-β-R monoclonal antibody[ies are] is CBE11 and at least one anti- LT-β-R monoclonal antibody is BHA10.
- 47. The pharmaceutical composition according to claim 41, wherein at least one [the] anti- LT-β-R monoclonal antibody[ies are] is CBE11 and at least one anti- LT-β-R monoclonal antibody is CDH10.

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The pharmaceutical composition according to claim 41, wherein <u>at least one</u> [the] anti- LT-β-R monoclonal antibody[ies are] <u>is</u> AGH1 and <u>at least one anti- LT-β-R monoclonal antibody is</u> CDH10.

49. The pharmaceutical composition according to any one of the claims 41-48 further comprising IFN-γ [as one of the LT-β-R activating agents].

Please add the following new claims to the application.

- 61. The method according to claim 7, wherein the anti- LT-β-R antibody has the same epitope specificity as an antibody produced by cell line CB.E11.1, ATCC accession number HB11793.
- 62. The method according to claim 7, wherein the anti- LT-β-R antibody has the same epitope specificity as an antibody produced by cell line BH.A10, ATCC accession number HB11795.
- 63. The method according to claim 9, wherein at least one anti- LT-β-R antibody has the same epitope specificity as an antibody produced by cell line CB.E11.1,

 ATCC accession number HB11793
- 64. The method according to claim 9, wherein at least one anti- LT-β-R antibody has the same epitope specificity as an antibody produced by cell line BH.A10, ATCC accession number HB11795.
- 65. The method according to claim 64, further comprising at least one anti- LT-β-R antibody having the same epitope specificity as an antibody produced by cell line CB.E11.1, ATCC accession number HB11793.

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- 66. The pharmaceutical composition according to claim 38, wherein the anti- LT-β-R antibody has the same epitope specificity as an antibody produced by cell line CB.E11.1, ATCC accession number HB11793.
- 67. The pharmaceutical composition according to claim 38, wherein the anti- LT-β-R antibody has the same epitope specificity as an antibody produced by cell line BH.A10, ATCC accession number HB11795.
- 68. The pharmaceutical composition according to claim 46, wherein at least one anti-LT-β-R antibody has the same epitope specificity as an antibody produced by cell line CB.E11.1, ATCC accession number HB11793.
- 69. The pharmaceutical composition according to claim 46, wherein at least one anti-LT-β-R antibody has the same epitope specificity as an antibody produced by cell line BH.A10, ATCC accession number HB11795.
- 70. The pharmaceutical composition according to claim 69, further comprising at least one anti- LT-β-R antibody having the same epitope specificity as an antibody produced by cell line CB.E11.1, ATCC accession number HB11793.

REMARKS

The instant application is a divisional of U.S.S.N. 08/875,560 ('560). In reply to the election requirement mailed October 14, 1999 in the '560 case Applicants elect prosecution of the method claims, species XI and XII (claims 7-17) and composition claims, species III and IV (claims 38-49) which are directed at compositions of at least two LT-β-R activating agents for treating neoplasia where at least one anti-LT-β-R activating agent is a LT-β-R antibody.

Upon entry of the present amendment, claims 7-17 and 38-49 will remain pending in the above-identified application as well as new claims 61-70.

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